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(54) Aluminum alloy and method for manufacturing aluminum-alloy member

(57) The invention offers an aluminum alloy that not only has high hardness accompanied by balanced ductility but also has high toughness and superior processability. The invention also offers a method for manufacturing an aluminum-alloy member that not only has high hardness accompanied by balanced ductility but also has high toughness and superior processability. The auminum alloy comprises (1) not less than 0.1 wt. % and not more than 8 wt. % Constituent A comprising one or more kinds of elements selected from the group consisting of titan Lm, vanadium, hafnium, and zirconium, 2 not less than 0.1 wt. % and not more than 20 wt. %

Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum, cerium, praseodymium, neodymium, mischmetal, calcium, strontium, and barium, and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium and lithium optionally further comprising not less than 0.1 wt.% and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium, molybdehum, silver, iron, cobalt, tantalum, and tungsten. In a second aspect the Al-alloy comprises the constituents D, B and C in the above specified amounts.

Description

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[0001] The present invention relates to an aluminum alloy and a method for manufacturing an aluminum alloy member and, more particularly, to an aluminum alloy combining good forgeability and high nardness and a method for manufacturing an aluminum-alloy member combining good forgeability and high hardness.

[0002] High-strength aluminum alloys have been in use in recent years that are produced by adopting a rapid solid-ification technique.

[0003] For instance, a published Japanese patent application *Tokukaihei* 1-275732 has disclosed that rapid solidification of a multi-element alloy expressed by a general formula Al_aM_bX_c produces a harocrystalline aluminum alloy having such mechanical properties as a tensile strength of 853 to 1.009 MPa, a yield strength of 804 to 941 MPa, and a hardness HV of 200 to 1,000. In the above formula Al_aM_bX_c, (1) "M" means one or more kinds of metal elements selected from the group consisting of chrome(Cr), manganese(Mn), iron(Fe), cobalt(Co₃, nickel(Ni), copper(Cu), zirconium(Zr), titanium(Ti), magnesium(Mg), and silicon(Si), (2) "X" means one or more kinds of metal elements selected from the group consisting of yttrium(Y), lanthanum(La), cerium(Ce), samarium(Sm), reodymium(No₃, niobium(Nb), and mischmetal(Mm), and (3) "a", "b", and "c" mean an atomic percent, "a" lying in the range of 50 to 95 atm. %, "b" in the range of 0.5 to 35 atm. %, and "c" 0.5 to 25 atm. %.

[0004] Another published Japanese patent application *Tokukaihei 6-184712* has disclosed an aluminum alloy having the composition expressed by a general formula Al_aLn_bM_c, where (1) "Ln" means one or more kinds of metal elements selected from the group consisting of mischmetal, yttrium, lanthanum, cerium, samanum, neodymium, hafnium, niobium, and tantalum, (2) "M" means one or more kinds of metal elements selected from the group consisting of vanadium, chrome, manganese, iron, cobalt, nickel, copper, zirconium, titanium, molybdenum, tungsten, calcium, lithium, magnesium, and silicon, and (3) "a", "b", and "c" mean an atomic percent, "a" lying in the range of 50 to 97.5 atm. %, "b" in the range of 0.5 to 30 atm. %, and "c" 0.5 to 30 atm. %. The aluminum alloy is a rapidly solidified aluminum alloy that has a cellular composite structure in which 5 to 50 vol. % amorphous phases surround nanocrystalline phases. The aluminum alloy is subjected to plastic working at a temperature higher than the crystallization temperature of the amorphous phase. Intermetallic compounds comprising two or more kinds of the above-described Ai. "Ln", and "M" are dispersed in the nanocrystalline matrix to form a structure having such mechanical properties as a tensile strength of 760 to 890 MPa and an elongation of 5.5 to 9.0%.

[0005] However, the aluminum alloy disclosed in the application *Tokukaihei 1-275732* has poor ductility and toughness, though it has very high tensile strength and hardness. Because this lack of sufficient ductility and toughness allows easy generation of cracks at the time of processing such as forging and upsetting, it is difficult to perform nearnet-shape forging with complicated shapes.

[0006] When forging is carried out by exploiting its superplasticity resulting from its nanocrystallinity, it is possible to impart complicated shapes. However, its poor ductility and toughness requires prolonged time for a single step of forging, causing a problem of reduced production efficiency, and hence an increase in manufacturing costs. Such a problem becomes serious when forming ornamental components that require complicated, fine shapes such as embossed letters on the surface.

[0007] Although the aluminum alloy disclosed in the application *Tokukaihei 6-184712* ensures a certain amount of ductility, it does not have sufficient mechanical properties to undergo near-net-shape forging with complicated shapes. In addition to that, because it uses material powders in which amorphous layers are formed, there is a problem of increased material cost.

[0008] The present invention is aimed at solving the above-described problems. An object of the present invention is to offer an aluminum alloy that not only has high hardness accompanied by balanced ductility but also has high toughness and superior processability.

[0009] Another object of the present invention is to offer a method for manufacturing an aluminum-alloy member that not only has high hardness accompanied by balanced ductility but also has high toughness and super or processability. [0010] The first aspect of the present invention offers an aluminum alloy that comprises (1) not less than 0.1 wt. % and not more than 8 wt. % Constituent A comprising one or more kinds of elements selected from the group consisting of titanium (Ti), vanadium (V), hafnium (Hf), and zirconium (Zr), (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of anthanum (La), cerium (Ce), praseodymium (Pr), neodymium (Nd), mischmetal (Mm), calcium (Ca), strontium (St), and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li).

[0011] The second aspect of the present invention offers another aluminum alloy that comprises 1) not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium (Nb), molybdenum (Mo), silver (Ag), iron (Fe), cobalt (Co), tantalum (Ta), and tungsten (W), (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum (La), cerium (Ce), praseodymium (Pr), neodymium (Nd), mischmetal (Mm),

calcium (Ca), strontium (St), and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li) [0012] The third aspect of the present invention offers a method for manufacturing an aluminum-alloy member made of the following aluminum alloy: The aluminum alloy comprises (1) not less than 0.1 wt. % and not more than 8 wt. % Constituent A comprising one or more kinds of elements selected from the group consisting of titanium (Ti), vanadium (V), hafnium (Hf), and zirconium (Zr), (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum (La), cerium (Ce), praseodymium (Pr, neodymium (Nd), mischmetal (Mm), calcium (Ca), strontium (St), and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li). First, a preform comprising the aluminum alloy is produced. Next, the preform is heated up to a temperature not lower than 200 °C and not higher than 600 °C at a temperature rising rate of not less than 2 °C/sec and not more than 200°C/sec. Then, the heated preform is subjected to hot-working.

[0013] The fourth aspect of the present invention offers a method for manufacturing an aluminum-alloy member made of the following aluminum alloy: The aluminum alloy comprises (1) not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium (Nb, molybdenum (Mo), silver (Ag), iron (Fe), cobalt (Co), tantalum (Ta), and tungsten (W). (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum (La), cerium (Ce), praseodymium (Pr), neodymium (Nd), mischmetal (Mm), calcium (Ca), strontium (St, and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li). First, a preform comprising the aluminum alloy is produced. Next, the preform is heated up to a temperature not lower than 200°C and not higher than 600 °C at a temperature rising rate of not less than 2°C/sec and not more than 200°C/sec. Then, the heates preform is subjected to hot-working.

[0014] The first to fourth aspects of the present invention offer an aluminum alloy that not only has high hardness accompanied by balanced ductility but also has high toughness and superior processability and a method for manufacturing an aluminum-alloy member that not only has high hardness accompanied by balanced ductility but also has high toughness and superior processability.

[0015] The first aspect of the present invention offers an aluminum alloy that comprises (1) not less than 0.1 wt. % and not more than 8 wt. % Constituent A comprising one or more kinds of elements selected from the group consisting of titanium (Ti), vanadium (V), hafnium (Hf), and zirconium (Zr), (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum (La), cerium (Ce), praseodymium (Pr), neodymium (Nd), mischmetal (Mm), calcium (Ca), strontium (St), and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li).

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[0016] Such a composition facilitates the formation of complicated shapes because it reduces the strength of the aluminum alloy in the temperature range for processing. This reduces the number of times of forming (forging) untitle last shape in comparison with the conventional products, and therefore reduces the processing cost.

[0017] This composition also increases the hardness of the aluminum alloy, and increased hardness suppresses the generation of surface flaws on members made of the aluminum alloy of the present invention during their manufacturing processes, reducing the fraction defective of the products.

[0018] The addition of a small amount of Ti, V, Hf, and Zr, which are used in Constituent A, can reduce the grain size of aluminum, increasing the hardness of the aluminum alloy. Intermetallic compounds between these elements and aluminum are deposited or crystallized out at the center of the individual crystal grains of aluminum (one place per crystal grain). If the content of Constituent A is less than 0.1 wt. %, the above-mentioned effect of increased hardness cannot be obtained. If the content of Constituent A is more than 8 wt. %, although the hardness of the aluminum allourincreases, the ductility, critical upsetting ratio, and other properties decrease, making it difficult to perform near-net-shape forging with complicated shapes, resulting in the reduction in forgeability.

[0019] The above-mentioned upsetting ratio is expressed in $(L0 - L1)/L0 \times 100$ (%), where L0 is the sample length in the upsetting direction before the upsetting work, and L1 after the upsetting work. The critical upsetting ratio is defined as the upsetting ratio at which cracks begin to develop at the periphery of the workpiece when upsetting is performed at a forging rate of 0.5 mm/sec. If the critical upsetting ratio is 70% or more, the sample is considered to have sufficient forgeability.

[0020] The elements La, Ce, Pr, Nd, Mm, Ca, Sr, and Ba, which are used in Constituent B, have an effect that a small amount of their addition can deposit a large amount of intermetallic compounds having high hardness. The deposition of intermetallic compounds increases the hardness of the aluminum alloy. The intermetallic compounds between these elements and aluminum are deposited or crystallized out at grain boundaries of aluminum. If the content of Constituent B is less than 0.1 wt. %, the above-mentioned effect cannot be obtained. If the content of Constituent B is more than 20 wt. %, although the hardness of the aluminum alloy increases, the ductility and other properties

deteriorate, reducing the forgeability.

[0021] The elements Mg and Li, which are used in Constituent C, have an effect that they can increase the hardness of the aluminum alloy when they are rapidly solidified in α -aluminum to form a supersaturated solid solution. If the content of Constituent C is less than 0.1 wt. %, the above-mentioned effect cannot be obtained. If the content of Constituent C is more than 20 wt. %, although the hardness of the aluminum alloy increases, the ductility, critical upsetting ratio, and other properties deteriorate, reducing the forgeability.

[0022] When Constituents A, B, and C are added with the specified contents as shown above, because the aluminum having Constituent C as a solid solution has fine crystal grains and because the intermetallic compounds are deposited or crystallized out at grain boundaries, a structure is formed that has less tendency to overgrow with temperature. The formation of this structure enables the production of an aluminum alloy with superior balance between the hardness and forgeability.

[0023] If any one of Constituents A, B, and C lies beyond the specified range of content, the balance between the hardness and forgeability is destroyed, producing high hardness with low forgeability or high forgeability with low hardness.

[0024] When an aluminum alloy having the above-described structure is hot-worked and then its surface is polished by buffing or other means, the surface of the member made of this hot-worked aluminum alloy can easily obtain metallic luster.

[0025] In the aluminum alloy of the first aspect of the present invention, it is more desirable that the content of Constituent C be more than 5 wt. % and not more than 20 wt. %.

[0026] This content range, when the surface of the aluminum alloy is anodized to form an anodic oxide coating, enables the anodic oxide coating to obtain a shade of relatively low brightness such as brown or dark gray.

[0027] The shade of the anodic oxide coating can be changed by adjusting the kind and content of elements used in Constituent C and other Constituents.

[0028] The aluminum alloy of the first aspect of the present invention may further comprise not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium (Nb), molybdenum (Mo), silver (Ag), iron (Fe), cobalt (Co), tantalum (Ta), and tungsten (W).

[0029] This can offer an aluminum alloy having good forgeablity and higher hardness.

[0030] The elements Nb, Mo, Ag, Fe, Co, Ta, and W, which are used in Constituent D, have an effect that they can not only reduce the grain size of aluminum but also deposit a large amount of intermetallic compounds. As a result, the hardness of the aluminum alloy can be further increased. In this case, the intermetallic compounds are deposited or crystallized out at a plurality of places inside the individual crystal grains of the aluminum.

[0031] If the content of Constituent D is less than 0.1 wt. %, the above-mentioned effect cannot be obtained. If the content of Constituent D is more than 5 wt. %, although the hardness of the aluminum alloy increases, the ductility, critical upsetting ratio, and other properties deteriorate, reducing the forgeability.

[0032] In the aluminum alloy of the first aspect of the present invention, it is more desirable that Constituent A be Zr, Constituent B be Mm, and Constituent C be Mg. In this case, it is more desirable that the content of Constituent A be not less than 0.1 wt. % and not more than 3 wt. % and the content of Constituent B be not less than 0.1 wt. % and not more than 15 wt. %.

[0033] The respective use of Zr, Mm, and Mg as Constituents A, B, and C can offer an aluminum alloy with further enhanced balance between the hardness and forgeability.

[0034] The second aspect of the present invention offers another aluminum alloy that comprises (1) not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium (Nb), molybdenum (Mo), solver (Ag), iron (Fe), cobalt (Co), tantalum (Ta), and tungsten (W), (2) not less than 0.1 wt. % and not more than 20 wt. For Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum (La), cerum (Ce), praseodymium (Pr), neodymium (Nd), mischmetal (Mm), calcium (Ca), strontium (St), and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li), [0035] Such a composition facilitates the formation of complicated shapes because it reduces the strength of the aluminum alloy in the temperature range for processing. This reduces the number of times of forming (forging) until the last shape in comparison with the conventional products, and therefore reduces the processing cost.

[0036] This composition also increases the hardness of the aluminum alloy. Increased hardness suppresses the generation of surface flaws on members made of the aluminum alloy of the present invention during their manufacturing processes, reducing the fraction defective of the products.

[0037] The elements Nb, Mo, Ag, Fe, Co. Ta. and W, which are used in Constituent D, have an effect that they can not only reduce the grain size of aluminum but also deposit a large amount of intermetallic compounds. As a result, the hardness of the aluminum alloy can be further increased. The intermetallic compounds produced by Constituent D are deposited or crystallized out at a plurality of places inside the individual crystal grains of the aluminum. If the content of Constituent D is less than 0.1 wt. 52, the above-mentioned effect cannot be obtained. If the content of

Constituent D is more than 5 wt. %, although the hardness of the aluminum alloy increases, the ductility, critical upsetting ratio, and other properties beteriorate, reducing the forgeability.

[0038] The elements La Ce, Pr, Nd, Mm, Ca, Sr, and Ba, which are used in Constituent B, have an effect that a small amount of their accion can deposit a large amount of intermetallic compounds having high hardness. The deposition of intermetallic compounds increases the hardness of the aluminum alloy. The intermetallic compounds produced by Constituent E are deposited or crystallized out at grain boundaries of aluminum.

[0039] If the content of Constituent B is less than 0.1 wt. %, the above-mentioned effect cannot be obtained. If the content of Constituent B is more than 20 wt. %, although the hardness of the aluminum alloy increases, the ductility, critical upsetting ratio, and other properties deteriorate, reducing the forgeability

[0040] The elements Mg and Li, which are used in Constituent C, have an effect that they can increase the hardness of the aluminum alloy when they are rapidly solidified in α-aluminum to form a supersaturated solid solution. If the content of Constituent C is less than 0.1 wt. %, the above-mentioned effect cannot be obtained. If the content of Constituent C is more than 20 wt. %, although the hardness of the aluminum alloy increases, the ductility, critical upsetting ratio, and other properties deteriorate, reducing the forgeability.

[0041] When Constituents D. B. and C are added with the specified contents as shown above, because the aluminum having Constituent C as a solid solution has fine crystal grains and because the intermetallic compounds are deposited or crystallized out at grain coundaries, a structure is formed that has less tendency to overgrow with temperature. The formation of this structure enables the production of an aluminum alloy with superior balance between the hardness and forgeability.

[0042] If any one of Constituents D, B, and C lies beyond the specified range of content, the balance between the hardness and forgeability is destroyed, producing high hardness with low forgeability or high forgeability with low hardness.

[0043] In the aluminum alloy of the second aspect of the present invention, it is more desirable that the content of Constituent C be more tran 5 wt. % and not more than 20 wt. %.

[0044] This content range, when the surface of the aluminum alloy is anodized to form an anodic oxide coating, enables the anodic oxide coating to obtain a shade of relatively low brightness such as brown or dark gray. The shade of the anodic oxide coating can be changed by adjusting the kind and content of elements used in Constituent C and other Constituents.

[0045] In the aluminum alloys of the first and second aspects of the present invention, it is more desirable that the aluminum alloys be further provided with an anodic oxide coating.

[0046] As mentioned acove, the shade of an anodic oxide coating can be changed by adjusting the kind and content of elements used in the individual Constituents. This enables the production of aluminum alloys provided with anodic oxide coatings having different shades. As a result, the painting process of the product can be omitted by using an anodic oxide coating having relatively high hardness as the protective coating of the aluminum alloy and by adjusting the shade of the anodic cxide coating so as to conform to the shade required in the product using the aluminum alloy. Consequently, the manufacturing cost of the product using the aluminum alloy can be reduced.

[0047] In the aluminum alloys of the first and second aspects of the present invention, it is more desirable that the anodic oxide coating have a lightness less than 50.

[0048] The lightness is measured by spectrophotometric colorimetry using a chromaticity meter (Japanese Industrial Standard JIS Z 8729; the _*a*b* color-expressing system). The light source for the measurement is D65 (the International Lighting Committee) the ISO standard light) with a color temperature of 6.504K.

[0049] In the aluminum alloys of the first and second aspects of the present invention, the anodic oxide coating may be formed on the surface of an aluminum-alloy base material. In this case, the base material may have an electrical conductivity less than 20% along (International Annealed Copper Standard).

[0050] The present in lentors have found that as the electrical conductivity of a base material decreases, the base-material element forms more solid solutions with the anodic oxide coating, giving a shade of relatively low brightness such as brown to the anodic oxide coating. The present inventors have also found that the base material requires to have an electrical conductivity less than 20 % IACS in order to give a shade of relatively low brightness such as brown to the anodic oxide coating.

[0051] In the aluminum alloys of the first and second aspects of the present invention, the anodic oxide coating may have a shade of brown lask gray, or dark brown.

[0052] When a compensation required to have a low-bright shade such as brown in the final product, the use of the aluminum alloy of the present invention makes it possible to obtain the required shade by adjusting the kind and content of elements used in the individual Constituents. This simplifies the traditionally required painting process of the component. Consequently, the manufacturing cost of the component can be reduced.

[0053] The aluminum aloys of the first and second aspects of the present invention may have aluminum crystals and intermetallic compaints. In this case, the aluminum crystals may have an average grain diameter of 1,000 nm or less and the intermetal a compounds may have an average grain diameter of 500 nm or less.

[0054] This enables the aluminum alloy to obtain high forgeability without losing the high hardness.

[0055] If the aluminum crystals have an average grain diameter more than 1,000 nm or the intermetallic compounds have an average grain diameter more than 500 nm, although the aluminum alloy improves its forgeability by improving its ductility, critical upsetting ratio, and other properties, it decreases its hardness.

[0056] In the aluminum alloys of the first and second aspects of the present invention, it is more desirable that the aluminum crystals have an average grain diameter of 500 nm or less and that the intermetallic compounds have an average grain diameter of 300 nm or less.

[0057] This enables the aluminum alloy to obtain higher hardness without losing its forgeability such as ductity and critical upsetting ratio when higher hardness is required.

[0058] The aluminum alloys of the first and second aspects of the present invention may have a Rockwell B hardness (H_{RB}) not less than 50 and not more than 100. In this case, the aluminum alloy may have a critical upsetting ratio of 70% or more at temperatures not lower than 200 °C and not higher than 600 °C and an elongation of 10% or more at 20 °C.

[0059] The hardness H_{RB} not less than 50 and not more than 100 means sufficiently high hardness in comparison with the conventional ingot aluminum alloys such as A5052. This high hardness suppresses the generation of surface flaws during the manufacturing process, thereby significantly reducing the ratio of defective products due to the surface flaws. If the hardness H_{RB} is less than 50, as in the conventional ingot aluminum alloys, it is difficult to suppress the generation of surface flaws during the manufacturing process. If the hardness H_{RB} is more than 100, such properties as the elongation at 20 °C and critical upsetting ratio deteriorate, reducing the forgeability.

[0060] The use of an aluminum alloy having the above-described critical upsetting ratio and elongation allows one or two processes of hot-working at temperatures not lower than 200 °C and not higher than 600 °C, facilitating the near-net-shape forging of components with complicated shapes. If the aluminum alloy has a critical upsetting ratio less than 70% at temperatures not lower than 200 °C and not higher than 600 °C or an elongation less than 10% at room temperature (20 °C), one or two processes of hot-working (near-net-shape forging) for obtaining components with complicated shapes generates work cracking of the components during the forging.

[0061] It is more desirable that the aluminum alloy of the first aspect of the present invention comprises (1; not less than 1.5 wt. % and not more than 2.5 wt. % Constituent A. (2) not less than 3 wt. % and not more than 6 wt. % Constituent B, (3) not less than 4 wt. % and not more than 6 wt. % Constituent C, and (4) not less than 1 wt. % and not more than 1.5 wt. % Constituent D.

[0062] The above-mentioned selection of the content ranges of Constituents A. B. C, and D enables the aluminum alloy to obtain a more enhanced balance between the hardness and workability (forgeability).

[0063] It is more desirable that the aluminum alloy of the second aspect of the present invention comprises (1) not less than 1.5 wt. % and not more than 2.5 wt. % Constituent D, (2) not less than 3 wt. % and not more than 6 wt. % Constituent B, and (3) not less than 4 wt. % and not more than 6 wt. % Constituent C.

[0064] The above-mentioned selection of the content ranges of Constituents D, B, and C enables the aluminum alloy to obtain a more enhanced balance between the hardness and workability (forgeability).

[0065] The third aspect of the present invention offers a method for manufacturing an aluminum-alloy member made of the following aluminum alloy: The aluminum alloy comprises (1) not less than 0.1 wt. % and not more than 8 wt. % Constituent A comprising one or more kinds of elements selected from the group consisting of titanium (Ti), variadium (V), hafnium (Hf), and zirconium (Zr), (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum (La), cerium (Ce), praseodym.um (Pr), neodymium (Nd), mischmetal (Mm), calcium (Ca), stront.um (St), and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li). First, a preform comprising the aluminum alloy is produced. Next, the preform is heated up to a temperature not lower than 200°C and not higher than 600°C at a temperature rising rate of not less than 2°C/sec and not more than 200°C/sec. Then, the heated preform is subjected to hot-working.

[0066] This procedure enables the easy production of an aluminum-alloy member having high hardness and a complicated shape notwithstanding the considerably reduced number of times of working during the hot-working process in comparison with the conventional methods.

[0067] If the temperature during the heating process idegasification process) of the preform is higher than 600 °C or the temperature-rising rate is less than 2 °C /sec or more than 200°C sec, the hot-working produces an aluminum alloy with sec or more than 200°C/sec, the hot-working produces an aluminum alloy with reduced hardness resulting from the coarsened grains of aluminum crystals and intermetallic compounds. If the heating temperature of the preform is lower than 200 °C, it is difficult to give the preform sufficient strength because of the insufficient bonding between the grains constituting the preform. This reduces the critical upsetting ratio at temperatures not lower than 200 °C and not higher than 600 °C and an elongation at room temperature (20 °C), deteriorating the forgeability.

[0068] In the method for manufacturing an aluminum-alloy member in the third aspect of the present invention, the aluminum alloy may further comprise not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one

or more kinds of elements selected from the group consisting of niobium (Nb), molybdenum (Mo), silver (Ag), iron (Fe), cobalt (Co), tantalum (Ta), and tangsten (W).

[0069] The fourth aspect of the present invention offers a method for manufacturing an aluminum-alloy member made of the following aluminum alloy: The aluminum alloy comprises (1) not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium (Nb), molybdenum (Mo), silver (Ag), iron (Fe), cobalt (Co), tantalum (Ta), and tungsten (W), (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum (La), cerium (Ce), praseodymium (Pr), neodymium (Nd), mischmetal (Mm), calcium (Ca), strontium (St), and barium (Ba), and (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium (Mg) and lithium (Li). First, a preform comprising the aluminum alloy is produced. Next, the preform is heated up to a temperature not lower than 200°C and not higher than 600 °C at a temperature using rate of not less than 2°C/sec and not more than 200°C/sec. Then, the heated preform is subjected to hot-working.

[0070] This procedure enables the easy production of an aluminum-alloy member having high hardness and a complicated shape notwithstanding the considerably reduced number of times of working during the hot-working process in comparison with the conventional methods.

[0071] If the heating temperature of the preform is higher than 600 °C or the temperature-rising rate is less than 2 °C/sec or more than 200°C/sec, the hot working produces an aluminum alloy with reduced hardness resulting from the coarsened grains of aluminum crystals and intermetallic compounds. If the heating temperature of the preform is lower than 200 °C, the preform becomes brittle because of the insufficient bonding between the grains constituting the preform. This reduces the critical upsetting ratio at temperatures not lower than 200 °C and not higher than 600 °C and an elongation at room temperature (20 °C), deteriorating the forgeability.

[0072] In the methods for manufacturing aluminum-alloy members in the third and fourth aspects of the present invention, it is more desirable that the heating temperature of the preform be not lower than 350 °C and not higher than 450 °C.

[0073] The above-mentioned selection of the heating temperature enables the aluminum-alloy member to easily obtain a more enhanced balance between the hardness and forgeability.

[0074] In the methods for manufacturing aluminum-alloy members in the third and fourth aspects of the present invention, it is desirable that the die temperature for the hot-working be about 400 °C.

[0075] In the methods for manufacturing aluminum-alloy members in the third and fourth aspects of the present invention, the step for producing the preform may include a step for forming rapidly solidified powders of aluminum alloy.

[0076] In the methods for manufacturing aluminum-alloy members in the third and fourth aspects of the present invention, the step for producing the preform may employ the OSPREY method.

[0077] In the methods for manufacturing aluminum-alloy members in the third and fourth aspects of the present invention, the step for producing the preform may include a step for forming powders produced by pulverizing rapidly solidified ribbons of aluminum alloy.

PREFERRED EMBODIMENTS OF THE INVENTION

40 [0078] The following is an explanation of the preferred embodiments of the present invention.

Embodiment 1

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[0079] Aluminum-alloy powders having a composition shown in the columns for Experimental Examples 1 to 11 in Table 1 were produced by using a gas-atomization device. In the gas-atomization method, a nitrogen gas was blown onto a molten aluminum alloy cropping from a nozzle having a hole 2 mm in diameter. The nitrogen gas was pressurized at 100 kgf/cm², in this case, air or an inert gas such as argon may be used in place of the nitrogen gas.

[0080] Powders of a 2014 a uninum alloy were also produced under the same condition of gas atomization as described above. The spacing between dendrite arms in the powder structure of the 2014 aluminum alloy was measured to estimate the cooling rate in the foregoing process. The result demonstrates that the production of powders having a particle diameter of 150 μ m corresponds to a cooling rate of 1.0 \times 10³ C/sec.

[0081] The aluminum-alloy powders were sieved out to obtain powders having a diameter less than 150 μ m. The obtained aluminum-alloy powders were press-formed to produce preforms. The preforms were heated up to temperatures of 350 to 400 °C at a temperature rising rate more than 2 °C/sec, actually at 10 °C/sec, as shown in Table 1 as the heating and degasifying treatments.

[0082] Subsequently, the preforms were inserted into a die kept at 400 °C to solidify the powders under a surface pressure of 9 t/cm². The fine structure and mechanical properties of the obtained solidified bodies were examined. The results are shown in Tables 1 and 2.

[0083] The grain diameters shown in Table 1 were determined by the following method: First, a section of a solidified body was mirror polished. Second, micrographs of the fine structure were taken by using a high-resolution scanning electron microscope (SEM) at 50,000 power. Finally, the individual micrographs were input into a personal computer to process the pictures for measuring the grain diameters of the aluminum crystals and intermetallic compounds. Because the aluminum crystals and intermetallic compounds have a different contrast on the micrograph, they are easily distinguished. The grain diameters were measured on three visual fields of each Experimental Example. Table 1 shows the average value of the measured results.

Table 1

					•
10	Company of the second s	Preform-heati	ng condition	Grain diameter of	Grain diameter of
Lowaer-		Illimate Temperature.	Temperature.	aluminum	
Solidined	(%)	temperature	rising rate	crystals	compounds
poq	(W.E. 70)	5	(C/sec)	(mm)	(mm)
· ·	At 27 8Mm 18Mg	400	10	009	300
- · :	MO1 272 8Mm 10Mg	400	01	200	500
	A) off Askm QMr		10	500	200
6 T	A1 9Mo 3Mm 11Mo		10	650	400
-	A1 5V - VNb - 3Mm - 8Mg	350	10	700	300
	A) 67: - 3Mm - 973 - 6Mr		10	850	400
0	A) Or SMm - SMe		10	200	300
	A1 - 221 - OMIN - 16Mv	400	10	500	300
0	A1 37r 4 5Mm - 8Mg	400	10	500	300
0.01	A1 22r - 8Mm - 8Mr - 3Mo	400	10	200	300
	A1 97 8Mm 8Ma 3Ta	400	10	500	300
	W	400	0.5	1,200	700
71	A - 27r - 8Mm - 16Mg	180	10	300	200
1.0		650	10	2,000	009
	A) - 22r - 8Mm - 25Me		10	009	700
GT	A1 10% RMm 8Mg		10	000	009
2:1	A1 . 3Zr = 21Mm = 6Mg	400	01	500	750
31	Al 2Zr - 3Mm - 6Mg - 10Mo		10	700	900
:	Al - 10Th - 6Mm - 10Mg -	400	10	009	800
	ÎZ X				

Table 2

					Ch. d. of	Plactrical	Lightness of
Room- Room-	Room.		Upsetting	Critical		הוכרנו זכמו	clumito
rafuro	tempera	ture	tempera-	upsetting ratio	alumite	conductiv-	aramire
	i terretari		, or the	(%)		ıty	•
nardness clougatum		ŧ	£ (£)			(%IACS)	,
11 _{KB}	(8)		(2)	- X	Brown	12.6	22.5
	2 :	-	007	75	Dark grav	13.0	23.6
01 10	2 :		000	08	Brown	12.0	21.5
36			000	85	Dark grav	16.0	33.5
1	: ::::::::::::::::::::::::::::::::::::		000	75	Dark gray	17.0	30.5
68	2 3		900	80	Dark gray	9.0	20.5
35			400	88	Dark gray	18.2	45.8
			007	**	Brown	15.2	43.1
2 2	2; 2		400	11.	Brown	17.3	47.5
	2 =		400	36.	Dark gray	27	36.9
	3 2		400	81	Dark gray	11.6	33.2
:	2 2	:	400	90	Light gray	25.0	63.9
-		1	400	50	Dark gray	10.5	31.8
0.5	2 6		400	95	Light gray	24.0	65.1
	17.		400	45	Dark gray	10.0	21.6
	2 4		400	65	Dark gray	18.0	28.4
) 		400	67	Dark gray	15.0	27.3
2 3	; ;		500	89	Dark gray	12.0	32.4
9 9	9	!	500	7.1	Dark brown	14.0	26.5

[0084] The fine structures of the solidified bodies of Experimental Examples 1 to 11 were examined by the above-described method.

[0085] As is seen in Table 1, it was confirmed that all the Experimental Examples 1 to 11 have both aluminum crystals and intermetallic compounds. It was also confirmed that all the Experimental Examples 1 to 11 have aluminum crystals less than 1,000 nm in grain diameter and intermetallic compounds less than 500 nm in grain 5 ameter.

[0086] Next, aluminum-alloy powders having a composition shown in the columns for Experimental Examples 12 to 19 in Table 1 were produced by a method similar to that used for Experimental Examples 10 11 Experimental Examples 12 to 19 were produced by using these powders. The heating conditions for the preferms of Experimental Examples 12 to 19 are also shown in Table 1. The fine structures of Experimental Examples 12 to 19 were also examined by the same method as in Experimental Examples 1 to 11.

[0087] The following measurements were carried out on Experimental Examples 1 to 19 shown in Table 1; hardness HRB at room temperature (20 °C), tensile strength at room temperature, critical upsetting ratio, and the shade of an anodic oxide coating (alumite) and other properties when the anodic oxide coating was formed on the surface.

[0088] As shown in Table 2, all the Experimental Examples 1 to 11 have a room-temperature hardness H_{RS} more than 50 and less than 100, an elongation not less than 10%, and a critical upsetting ratio more than 70%.

[0089] The surfaces of the solid bodies of Experimental Examples 1 to 11 were anodized to form an anodic exide coating (alumite). The shade of the alumite was examined. As is seen in Table 2, the results showed that all the Experimental Examples 1 to 11 have a dark shade such as brown or dark gray. The lightness of the alumite was measured; the result showed that all the Experimental Examples 1 to 11 have a lightness less than 50. The electrical conductivity of the matrices of the solidified bodies of Experimental Examples 1 to 19 was measured. As is seen in Table 2, the result showed that when the electrical conductivity is less than 20 %IACS, the shade of the alumite is dark (less than 50 in lightness) such as brown. Incidentally, all the Experimental Examples 1 to 11 have an electrical conductivity less than 20 %IACS.

[0090] The fine structures and mechanical properties of Experimental Examples 12 to 19 are discussed in the following:

[0091] Experimental Example 12 has a room-temperature hardness as low as 49 as is seen in Table 2. This is attributable to the grain diameter of the aluminum crystals as large as 1,200 nm resulting from the temperature-nsing rate as low as 0.5 °C/sec in the preform-heating conditions as can be seen in Table 1.1' the room-temperature hardness is less than 50, surface flaws and other defects tend to be generated during the manufacturing process, causing a yield reduction as in the conventional products.

[0092] Experimental Example 13, although having a room-temperature hardness exceeding 100, has practically no elongation and a critical upsetting ratio as low as 50%. This is attributable to the fact that the ultimate temperature was 180 °C in the preform-heating conditions, i.e., the preform was not heated up to a temperature exceeding 200 °C.

[0093] Experimental Example 14 also has a room-temperature hardness as low as 46, as is seen in Table 2. This is attributable to the grain diameter of the aluminum crystals as large as 2,000 nm, which is more than necessary, resulting from the ultimate temperature as high as 650 °C in the preform-heating conditions as can be seen in Table 1.

[0094] Experimental Example 15 contains, in its composition, more Mg than the content specified for the aluminum alloy of the present invention as is seen in Table 1. Consequently, as shown in Table 2, a though sufficiently high in room-temperature hardness. Experimental Example 15 has low elongation and critical upsetting ratio, and hence low forgeability.

[0095] Experimental Example 16 contains more Zr than the content specified for the aium num alloy of the present invention as is seen in Table 1. Consequently, as shown in Table 2, although sufficiently high in room-temperature hardness, Experimental Example 16 has low elongation and critical upsetting ratio.

[0096] Experimental Example 17 contains more Mm than the content specified for the alumnum alloy of the present invention as is seen in Table 1. Consequently, as shown in Table 2, although sufficiently high in room-temperature hardness, Experimental Example 17 has low elongation and critical upsetting ratio.

[0097] Experimental Example 18 contains more Mo than the content specified for the alumnum alloy of the present invention as is seen in Table 1. Consequently, as shown in Table 2, although sufficiently high in room-temperature hardness, Experimental Example 18 has low elongation and critical upsetting ratio.

[0098] Experimental Example 19 contains more Ti and Nb than the contents specified for the aluminum alloy of the present invention as is seen in Table 1. Consequently, as shown in Table 2, although sufficiently high in room-temperature hardness. Experimental Example 19 has low elongation.

[0099] The anodizing was carried out by the following process: First, the surface of a solid fed body was cut. Second, the solidified body subjected to the cutting work was cleaned by caustic soda. Finally anodizing was conducted up to a coating thickness of about $10 \, \mu m$.

[0100] The structure in the vicinity of the boundary between the anodic oxide coating and case material (matrix) was examined on the individual Experimental Examples 1 to 19 by using a high-resolution scanning electron microscope. The result was that the reflected electron image of the structure demonstrates the existence of intermetallic compounds

in the anodic oxide coating. When the shade of an anodic oxide coating (alumite) becomes brown or dark gray, the alumite has an increased amount of the intermetallic compounds to a certain extent. More specifically, the intermetallic compounds occupy more than 20% of the area of the alumite.

[0101] It was also confirmed that when a sample made of the aluminum alloy of the present invention is ubset at high temperature and then its surface is polished by buffing or another simple means, the surface of the sample can easily obtain metallic luster.

Embodiment 2

[0102] Aluminum-alloy powders having a composition shown in the columns for Experimental Examples 20 to 27 in Table 3 were produced by a method similar to that used for Embodiment 1 of the present invention. Experimental Examples 20 to 27 were produced by using these powders. Samples of solidified bodies were formed by a method basically similar to that used for Embodiment 1 of the present invention. The heating conditions for the preferms are shown in Table 3. The fine structures and mechanical properties of the solidified bodies were examined by a method similar to that used for Embodiment 1 of the present invention. The results are shown in Tables 3 and 4.

																					_
5		Grain diameter of	intermetanic	(mu)	400	300	300	300	250	300	300	500	009	300	700	700	000	1,000	000	000	006
15		neter of	E	crystals (nm)	650	700	400	500	400	200	200	400	1,200	500	2,000	200	009	200		1,000	1,200
25	Table 3	Preform-heating condition	Temperature-	rising rate (C/sec)	10	10	10	10	10	10	10	10	0.5	10	10	91	10	10		10	10
30	Tal	Preform-hea	Ultimate	temperature	400	350	400	400	400	350	400	400	400	180	650	400	400	400		400	400
35						NI, 2Mm 3Mg	3Mr	5Mm - 3Mo	J. OMG	Mm - 9Mc - 3Mo	Mm - 2Mc - 3W	13 5Mm - 21 i		3Mm -: 3Mg	9	- 25M.	Mm MM	. 10Mm - 11La -		m ~ 3Mg	1 - 4Mg
45		Composition		(wt. %)	AAC AAC	- 01A17	Al 1975 13Mm		A1 2A2 13Mr	900	- MWC - VC - VA		W 100 - 10			MA SMC IN	N - ZIMO - OMIT	VI = 3Zr = 10	ME	N - 10Mo - 3M	AI - 10W - 2Mm - 4Mg
50		vder-	diffed	<u>~</u>		} 	_: {	!		7	:				i		1	25			35

Table 4

'owder	Room	Room	Upsetting	Critical up	up. Shade of	Electrical	Lightness of
5	temperature	temperature	tempera-	setting ra	ra- alumite	conductiv-	alumite
=======================================	hardness	elongation	ture	tio		ity	
	1-1	(%)	ξ)	<u>(%</u>		(%IACS)	*1
	88		500	65	Light gray	24.3	65.2
	78	13	300	76	Light gray	26.8	8.69
<u>i</u>	7.4	13	400	81	Light yellow	26.5	68.2
!	7.3	20	400	36	Light yellow	24.0	61.4
:	84	14	400	85	Light yellow	24.5	59.8
!	88	12	200	92	Light gray	20.0	68.3
	87		200	85	Light gray	21.8	69.2
	85	10	400	79	Dark gray	18.9	36.3
:	44	25	400	96	Light gray	30.0	60.2
<u>i</u>	101	2	400	50	Dark gray	10.6	20.4
	48	22	400	95	Light gray	32.0	59.8
<u> </u>	112	0	400	45	Dark gray	9.0	15.8
	16	3	400	55	Dark gray	11.0	20.6
<u> </u>	91		400	45	Dark gray	8.0	17.8
<u> </u>	81	L	400	80	Dark gray	18.6	45.6
<u> 1</u> .	82	x	400	92	Dark gray	19.2	48.9

[0103] As is seen in Tables 3 and 4, the measured items for Experimental Examples 20 to 27 are the same as those for Embodiment 1 of the present invention. The measured results for all the items of Experimental Examples 20 to 27 are within the range specified for the aluminum aloy of the present invention. The surfaces of the samples were anodized similarly to Embodiment 1 of the present invention to form an anodic oxide coating (alumite). The shade and lightness of the alumite were examined. The electrical conductivity of the base material was also measured. As can be seen in Tables 3 and 4, the shade of the aluminum alloy can be changed to one such as dark gray or light yellow by adjusting the composition of the aluminum alloy.

[0104] Aluminum-alloy powders having a composition shown in the columns for Experimental Examples 28 to 35 in Table 3 were produced similarly to Experimental Examples 20 to 27. The powders were used to produce solidified bodies under the heating conditions for the preferms shown in Table 3. The fine structures and mechanical properties of the solidified bodies were examined similarly to Experimental Examples 20 to 27.

[0105] Experimental Example 28 was prepared by a temperature-rising rate lower than 2 °C/sec. Experimental Example 30 was prepared at an ultimate temperature higher than 600 °C. As a result, Experimental Examples 28 and 30 have aluminum-crystal grains and intermetal' c compounds both larger in diameter than the values desirable for the aluminum alloy of the present invention. Consequently, both Examples have a considerably low rocm-temperature hardness as shown in Table 4.

[0106] Experimental Example 29 was prepared at an ultimate temperature lower than 200°C. Consequently, although high in room-temperature hardness, Experimental Example 29 has low elongation and critical upsetting ratio.

[0107] Experimental Example 31 contains Constituent C of which Mg exceeds 20 wt. % in content. Consequently, although sufficiently high in room-temperature hardness, Experimental Example 31 has considerably low elongation and critical upsetting ratio.

[0108] Experimental Example 32 contains Constituent A of which the sum of Ti and V exceeds 8 wt. % in content. As a result. Experimental Example 32 has intermetallic compounds not only increased in the amount of deposition but also excessively grown. Consequently, although sufficiently high in room-temperature hardness, Experimental Example 32 has considerably low elongation and critical apsetting ratio.

[0109] Experimental Example 33 contains Constituent B of which the sum of Mm and La exceeds 20 wt. % in content. As a result. Experimental Example 33 also has intermetallic compounds not only increased in the amount of deposition but also excessively grown. Consequently, although sufficiently high in room-temperature hardness, Experimental Example 33 has considerably low elongation and critical upsetting ratio.

[0110] Experimental Example 34 contains more Mo than 5 wt. %. In this case also, the intermetallic compounds are excessively grown. Consequently, although the room-temperature hardness becomes high, the elongation decreases.
[0111] Experimental Example 35 contains more W than 5 wt. %. In this case also, the intermetallic compounds are not only increased in the amount of deposition but also excessively grown. Consequently, although the room-temperature hardness increases to a certain extent, the elongation decreases.

In [0112] As described above, the aluminum allay of the present invention has high hardness and good elongation and critical upsetting ratio (forgeability). It is also possible to obtain a member having metal luster by giving it simple polishing after hot-working.

[0113] Because the shade of the alumite can be changed by adjusting additive elements, a highly hard anodic oxide coating can be used not only as a protective spating but also as a colored layer that gives necessary coloring.

[0114] The aluminum alloy of the present invention can be used both as an exterior component of electronic devices, for example, and as a component of household electrical appliances, ornamental objects, cars, and other objects.

[0115] The present embodiments are to be considered in all respects as illustrative and not restrictive. The scope of the present invention is indicated by the appended claims rather than by the above-described embodiments. All changes that come within the meaning and range of equivalency of the claims are therefore intended to be embraced by the claims.

Claims

- 1. An aluminum alloy comprising:
 - (1) not less than 0.1 wt. % and not more than 8 wt. % Constituent A comprising one or more κ rds of elements selected from the group consisting of transium, variadium, hafnium, and zirconium.
 - (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more ands of elements selected from the group consisting of anthanum, cerium, praseodymium, neodymium, mischmetal, calcium, strontium, and barium, and
 - (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more ands of elements selected from the group consisting of magnesium and lithium.

- 2. An aluminum alloy as defined in claim 1, wherein Constituent C constitutes not less than 5 wt. % and not more than 20 wt. %.
- An aluminum alloy as defined in claim 1, the aluminum alloy further comprising not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of nilobium, molybdenum, silver, iron, cobalt, tantalum, and tungsten.
 - 4. An aluminum alloy as defined in claim 2, wherein:

Constituent A is zirconium,

Constituent B is mischmetal,

Constituent C is magnesium,

Constituent A constitutes not less than 0.1 wt. % and not more than 3 wt. %, and

Constituent B constitutes not less than 0.1 wt. % and not more than 15 wt. %.

5. An aluminum alloy comprising:

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- (1) not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium, molybdenum, silver iron, cobalt, tantalum, and tungsten,
- (2) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum, cerium, praseocymium, neodymium, mischmetal, calcium strontium, and barium, and
- (3) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium and lithium.

An aluminum alloy as defined in claim 5, wherein Constituent C constitutes not less than 5 wt. % and not more than 20 wt. %.

- An aluminum alloy as defined in one of the claims 1 to 6, the aluminum alloy being further provided with an anodic oxide coating.
 - 8. An aluminum alloy as defined in claim 7, wherein the anodic oxide coating has a lightness less than 50.
 - 9. An aluminum alloy as defined in claim 8, wherein:

the anodic oxide coating is formed on the surface of a base material made of the aluminum alloy, and the base material has an electrical conductivity less than 20 %IACS.

- **10.** An aluminum alloy as defined in claim 8, wherein the anodic oxide coating is a shade of brown, dark gray, or dark brown.
 - 11. An aluminum alloy as defined in one of the claims 1 to 6, the aluminum alloy having aluminum crystals and intermetallic compounds.

the aluminum crystals having an average grain diameter of 1,000 nm or less, the intermetallic compounds having an average grain diameter of 500 nm or less.

- 12. An aluminum alloy as defined in claim 11, wherein the aluminum crysta's have an average grain diameter of 500 nm or less and the intermetallic compounds have an average grain diameter of 300 nm or less.
- 13. An auminum alloy as defined in one of the claims 1 to 6, the aluminum alloy having:

a hardness H_{RB} not less than 50 and not more than 100. a critical upsetting ratio of 70% or more at temperatures not lower than 200 °C and not higher than 600 °C, and an elongation of 10% or more at 20 °C.

14. A method for manufacturing an aluminum-alloy member, the method comprising the steps of:

- (1) producing a preform comprising an aluminum alloy comprising:
 - (a) not less than 0.1 wt. % and not more than 8 wt. % Constituent A comprising one or more kinds of elements selected from the group consisting of titanium, vanadium, hafnium, and zirconium,
 - (b) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum, cerium, praseodymium, neodymium, mischmetal, calcium, strontium, and barium, and
 - (c) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium and lithium;
- (2) heating the preform up to a temperature not lower than 200 °C and not higher than 600 °C at a temperature rising rate of not less than 2°C/sec and not more than 200°C/sec; and
- (3) subjecting the heated preform to hot-working.

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- 15. A method for manufacturing an aluminum-alloy member as defined in claim 14, wherein the aluminum alloy further comprises not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium, molybdenum, silver, iron, cobalt, tantalum, and tungsten.
 - 16. A method for manufacturing an aluminum-alloy member, the method comprising the steps of:
 - (1) producing a preform comprising an aluminum alloy comprising:
 - (a) not less than 0.1 wt. % and not more than 5 wt. % Constituent D comprising one or more kinds of elements selected from the group consisting of niobium, molybdenum, silver, iron, cobalt, tantalum, and tungsten.
 - (b) not less than 0.1 wt. % and not more than 20 wt. % Constituent B comprising one or more kinds of elements selected from the group consisting of lanthanum, cerium, praseodymium, neodymium, mischmetal, calcium, strontium, and barium, and
 - (c) not less than 0.1 wt. % and not more than 20 wt. % Constituent C comprising one or more kinds of elements selected from the group consisting of magnesium and lithium;
 - (2) heating the preform up to a temperature not lower than 200 °C and not higher than 600 °C at a temperature rising rate of not less than 2°C/sec and not more than 200°C/sec; and
 - (3) subjecting the heated preform to hot-working.
 - 17. A method for manufacturing an aluminum-alloy member as defined in one of the claims 14 to 16, wherein the step of producing the preform includes a step of forming rapidly solidified powders of the aluminum alloy.
 - 18. A method for manufacturing an aluminum-alloy member as defined in one of the claims 14 to 16, wherein the step of producing the preform employs the OSPREY method.
 - 19. A method for manufacturing an aluminum-alloy member as defined in one of the claims 14 to 16, wherein the step of producing the preform includes a step of forming powders produced by pulverizing rapidly solidified ribbons of the aluminum alloy.



EUROPEAN SEARCH REPORT

Application Number EP 99 30 8574

tegory	Citation of document with indic of relevant passage		Rele to cla		CLASSIFICATION OF THE APPLICATION (Int.CI.7)
, ,X	PATENT ABSTRACTS OF J vol. 1999, no. 13, 30 November 1999 (199 & JP 11 236601 A (SUM LTD), 31 August 1999 * abstract *	9-11-30) ITOMO ELECTRIC IND	1,2,1		C22C21/00 C22C45/08 C22C21/06 B22F9/08 C25D11/04
x	EP 0 558 977 A (YOSHI 8 September 1993 (199 * page 3, line 26 - 1 * page 4, line 16 - 1 16,20; table 1 *	3-09-08)	1,5, 12.1	11, 7,19	
A	10,20, table 1		14,1	6	
X	PATENT ABSTRACTS OF C vol. 012, no. 117 (C- 13 April 1988 (1988-C -& JP 62 238345 A (Sk 19 October 1987 (1987 * abstract * * page 4; example 4;	487), 14-13) IOWA ALUM CORP), 7-10-19)	1,2,	4	TECHNICAL FIELDS SEARCHED (Int.Cl.7)
X	EP 0 610 006 A (TOYO 10 August 1994 (1994 * page 5, line 14 - table 3 *	-08-10)	1,4,	11,	C22C B22F
X	US 3 278 300 A (KICH 11 October 1966 (1960 * column 2, line 55 6-9; tables 11-16 *	5-10-11)	5		
X	FR 1 521 857 A (D. Co 26 July 1968 (1968-0 * page 2; table *		5		
		-/			
	The present search report has be				Example
	Page of search	28 February 2	1	i	impakis, E
X:pa Y:pa	CATEGORY OF CITED DOCUMENTS articularly relevant if taken alone uniquiarly relevant if combined with anothe current of the same category onlocing cell background	T 'theory or 3 E : earlier pare after the fir D : document L : document	nnoine uncert int document, ing cate sited in the ap itted for other	ying the but put prication reasons	nvention fished on, or



EUROPEAN SEARCH REPORT

Application Number EP 99 30 8574

	DOCUMENTS CONSIDER Citation of document with India		Relevant	CLASSIFICATION OF THE
ategory	of relevant passage		to claim	APPLICATION (Int.Cl.7)
4	EP 0 529 542 A (YOSHI	DA KOGYO KK)	1,5,11,	
`	3 March 1993 (1993-03		12.14.	
	3 March 1993 (1993-03			
	* page 3, line 6 - li		16,17,19	
A	EP 0 534 470 A (MASUM AKIHISA (JP); HIGASH 31 March 1993 (1993-0 * page 3, line 37 - p * page 4, line 31 -	KENJI (JP); ÝOSHIDA) 03-31) page 4, line 5 *	1,5,11, 12,14, 16,17,19	
Α	PATENT ABSTRACTS OF vol. 014, no. 115 (C) 5 March 1990 (1990-0) -& JP 01 316433 A (F) 21 December 1989 (1994 abstract * * example 10; table	-0696), 3-05) JRUKAWA ALUM CO LTD), 39-12-21)	5,17,19	
Α	PATENT ABSTRACTS OF	1APAN	7,8,10	
A	1	_	7,0,10	TECHNICAL FIELDS
	vol. 014, no. 037 (C		ì	SEARCHED (Int.CI.7)
	24 January 1990 (199	0-01-24)	ļ	
	-& JP 01 272740 A (S	HOWA ALUM CORP),	1	
	31 October 1989 (198		İ	
	* abstract *	,	1	1
	- abstract +		1	
A	PATENT ABSTRACTS OF vol. 014, no. 037 (C 24 January 1990 (199 -& JP 01 272739 A (S 31 October 1989 (198	-680), 0-01-24) HOWA ALUM CORP),	7,8,10	
	* abstract *	-	1	1
	-			
				1
				1
	1		1	
	1		}	
1	i		ļ	
	1		_i	
	The present search report has b	een drawn up for all claims		
	Pace or scarch	Date or ocmpetion of the search	1 .	Example:
	MUNICH	28 February 2000	1 1 4	limpakis, E
	HOUTCH	Zo repruary 2000	<u> </u>	TIMPAKIS, E
[CATEGORY OF CITED DOCUMENTS	Tir theory or ashalo		
		E : earlier patent or	ocument, but put	
X:5	anicularly relevant if taken alone articularly relevant if combined with anoth	after the filing co er D : document cited	are 'in the application	ra .
'.c	ocument of the same category	L : document sited	tor other reason	S
A . 16	ectinological background	2 mambas of the		
	ion-writer disclosure iternediate cocument	& , Temper of the : document	Service Services	y, corresponding
F - "	no continu properties i			

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 99 30 8574

This annex itsis the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on.

The European Patent Office is n no way liable for these particulars which are merely given for the purpose of information.

28-02-2000

cite	atent document ed in search repor	n.	Publication date	Patent family member(s)	Publication date
JP	11236601	A	31-08-1999	NONE	
EP	0558977	A	08-09-1993	JP 2954775 B JP 5222491 A DE 69310954 D DE 69310954 T US 5647919 A	27-09-199 31-08-199 03-07-199 08-01-199 15-07-199
JP	62238345	A	19-10-1987	JP 2101206 C JP 6021304 B	22-10-199 23-03-199
EP	0610006	A	10-08-1994	JP 7145441 A CA 2114285 A DE 69413571 D DE 69413571 T US 5573608 A	06-06-199 28-07-199 05-11-199 01-04-199 12-11-199
US	3278300	A	11-10-1966	NONE	
FR	1521857	Α	26-07-1968	NONE	
EP	0529542	A	03-03-1993	JP 5051684 A DE 69209588 D DE 69209588 T US 5415709 A	02-03-199 09-05-199 21-11-199 16-05-199
EP	0534470	A	31-03-1993	DE 69220164 D DE 69220164 T JP 2865499 B JP 6017178 A US 5405462 A US 5332456 A	10-07-199 08-01-199 08-03-199 25-01-199 11-04-199 26-07-199
JP	01316433	Α	21-12-1989	NONE	
JP	01272740	A	31-10-1989	NONE	
	01272739	Α	31-10-1989	NONE	

For more details about this annex ; see Official Journal of the European Patent Office, No. 12/82